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NEWS
                 Web Page for STN Seminar Schedule - N. America
NEWS 2 AUG 10 Time limit for inactive STN sessions doubles to 40
                 minutes
NEWS 3 AUG 18
                 COMPENDEX indexing changed for the Corporate Source
                 (CS) field
NEWS
     4 AUG 24
                 ENCOMPLIT/ENCOMPLIT2 reloaded and enhanced
NEWS
         AUG 24
                 CA/CAplus enhanced with legal status information for
                 U.S. patents
        SEP 09
                 50 Millionth Unique Chemical Substance Recorded in
                 CAS REGISTRY
NEWS 7 SEP 11
                 WPIDS, WPINDEX, and WPIX now include Japanese FTERM
                 thesaurus
NEWS 8 OCT 21 Derwent World Patents Index Coverage of Indian and
                 Taiwanese Content Expanded
NEWS 9
        OCT 21 Derwent World Patents Index enhanced with human
                 translated claims for Chinese Applications and
                 Utility Models
NEWS 10 OCT 27
                 Free display of legal status information in CA/CAplus,
                 USPATFULL, and USPAT2 in the month of November.
NEWS 11 NOV 23 Addition of SCAN format to selected STN databases
NEWS 12 NOV 23 Annual Reload of IFI Databases
NEWS EXPRESS MAY 26 09 CURRENT WINDOWS VERSION IS V8.4,
             AND CURRENT DISCOVER FILE IS DATED 06 APRIL 2009.
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FILE 'HOME' ENTERED AT 17:55:20 ON 23 NOV 2009

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SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 0.22

0.22

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STRUCTURE FILE UPDATES: 22 NOV 2009 HIGHEST RN 1193309-59-9 DICTIONARY FILE UPDATES: 22 NOV 2009 HIGHEST RN 1193309-59-9

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```
12 14 15 17 18 19 20 22 23 ring nodes:
1 2 3 4 5 6 7 8 9 10 24 25 26 27 28 29 chain bonds:
1 2 3 4 5 6 7 8 9 10 24 25 26 27 28 29 chain bonds:
1 2 7-14 7-15 8-22 8-23 9-17 9-18 10-19 10-20 12-24 ring bonds:
1 -2 1 -6 2 -3 3 -4 4 -5 5 -6 5 -7 6 -10 7 -8 8 -9 9-10 24-25 24-29 25-26 26-27 27-28 28-29 exact/norm bonds:
5 -7 6 -10 7 -8 7 -14 7 -15 8 -9 8 -22 8 -23 9 -10 9 -17 9 -18 10 -19 10 -20 12 -24 24 -25 24 -29 25 -26 26 -27 27 -28 28 -29 normalized bonds:
1 -2 1 -6 2 -3 3 -4 4 -5 5 -6 isolated ring systems: containing 24 :
```

G1:H,N

G2:C, H

G3:C,N

Match level :

chain nodes :

12:CLASS 14:CLASS 15:CLASS 17:CLASS 18:CLASS 19:CLASS 20:CLASS 22:CLASS 23:CLASS 25:CLASS 25:

L1 STRUCTURE UPLOADED

=> s 11 sss

SAMPLE SEARCH INITIATED 17:56:05 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 119757 TO ITERATE

1.7% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **INCOMPLETE** BATCH **COMPLETE** PROJECTED ITERATIONS: 2374598 TO 2415682

PROJECTED ANSWERS:

1.2 0 SEA SSS SAM L1

=> s 11 full

THE ESTIMATED SEARCH COST FOR FILE 'REGISTRY' IS 185.40 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:y FULL SEARCH INITIATED 17:56:10 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 2402134 TO ITERATE

0 TO

81.8% PROCESSED 1963985 ITERATIONS

190 ANSWERS 190 ANSWERS

0 ANSWERS

83.3% PROCESSED 2000000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.23

FULL FILE PROJECTIONS: ONLINE **INCOMPLETE** BATCH **COMPLETE** 2402134 TO 2402134 PROJECTED ITERATIONS: 190 TO 273

1.3 190 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS

SINCE FILE

FULL ESTIMATED COST

PROJECTED ANSWERS:

ENTRY SESSION 186.36 186.58

TOTAL

FILE 'CAPLUS' ENTERED AT 17:56:37 ON 23 NOV 2009 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

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FILE COVERS 1907 - 23 Nov 2009 VOL 151 ISS 22
FILE LAST UPDATED: 22 Nov 2009 (20091122/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Aug 2009
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

CAplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2009.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

During November, try the new LSUS format of legal status information in the CA/CAplus family databases for free! Complete details on the number of free displays and other databases participating in this offer appear in NEWS 10.

=> s 13 full L4 20 L3

=> d ibib abs hitstr tot THE ESTIMATED COST FOR THIS REQUEST IS 112.80 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:y

GΙ

L4 ANSWER 1 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2006:335893 CAPLUS

DOCUMENT NUMBER: 144:390943

TITLE: Preparation of arylpiperazine derivatives as tubulin inhibitors for treatment of proliferation or cancer

INVENTOR(S): Betzemeier, Bodo; Krist, Bernd; McConnell, Darryl; Steurer, Steffen; Impagnatiello, Maria;

Wever-Czernilofsky, Ulrike; Hilberg, Frank; Brueckner, Ralph; Daiimann, Georg; Heckel, Armin; Kley, Joerg;

Lehmann-Lintz, Thorsten; Roth, Gerald

PATENT ASSIGNEE(S): Boehringer Ingelheim International G.m.b.H., Germany SOURCE:

Eur. Pat. Appl., 55 pp.

CODEN: EPXXDW DOCUMENT TYPE: Patent

LANGUAGE . English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
EP 1645556	A1 20060412	EP 2004-23926	20041007
R: AT, BE, CH,	DE, DK, ES, FR, G	GB, GR, IT, LI, LU, NL,	SE, MC, PT,
IE, SI, LT,	LV, FI, RO, MK, C	CY, AL, TR, BG, CZ, EE,	HU, PL, SK, HR
PRIORITY APPLN. INFO.:			20041007
OTHER SOURCE(S):	CASREACT 144:3909	43; MARPAT 144:390943	

R10 Ŕ9 R3 R2 Ι Ċ1

The title arylpiperazine derivs. I [wherein A = mono- or bicyclic aryl; R1 and R2 = independently H, halo, CN, (un) substituted alkyl, alkoxy, etc.; R3 = H, halo, CN, alkyl, or alkoxy; or R2 and R3 = (un)substituted -O-(CH2)p-O- ring; R4 and R5 = independently H or alkyl; R6-R10 =

independently H, halo, NO2, CN, (un)substituted alkyl, NH2, alkoxy, etc.; X and Y = independently CH, CF, or N; n and p = independently 1 or 2], or pharmaceutically acceptable salts, derivs., tautomers, or solvates thereof were prepared as tubulin inhibitors for the treatment of proliferative diseases or cancer (no data). For example, 4-amino-3,5-dichlorobenzoic acid was reacted with 1-(3-chlorophenyl)-piperazine in DMF at 50 °C in the presence of TBTU to give II (47 %). The title compds. showed inhibitory activity with IC50 < 10 µM in vitro cytotoxicity assay. Formulations as tablets, coated tablets, capsules, or ampoules were described.

IΤ 882695-10-5P

> RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug candidate; preparation of arylpiperazine derivs. as tubulin inhibitors for treatment of proliferation or cancer)

RN 882695-10-5 CAPLUS

CN Methanone, [4-(3,5-dimethoxyphenyl)-1-piperazinyl](5,6,7,8-tetrahydro-1naphthalenvl) - (CA INDEX NAME)

OS.CITING REF COUNT:

- THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
- REFERENCE COUNT:
- 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

<12/04/2007> Erich Leese

1

L4 ANSWER 2 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2005:979643 CAPLUS

DOCUMENT NUMBER: 143:266686

TITLE: Preparation of tetralin derivatives as histamine H3

receptor antagonists
INVENTOR(S): Beavers, Lisa Selsam; Gadski, Robert Alan; Hipskind,

Philip Arthur; Jesudason, Cynthia Darshini; Lindsley,
Craig William; Lobb, Karen Lynn; Pickard, Richard Todd

PATENT ASSIGNEE(S): Eli Lilly and Company, USA

SOURCE: PCT Int. Appl., 49 pp.

OURCE: PCT Int. App1 CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA'	PATENT NO.			KIND DATE				APPL	ICAT	ION	NO.		D.	ATE				
	2005 2005						2005 2006			WO 2	005-	US54	91		2	0050	222	
	W:	CN, GE, LK,	CO, GH, LR,	CR, GM, LS,	CU, HR, LT,	CZ, HU, LU,	AU, DE, ID, LV, PL,	DK, IL, MA,	DM, IN, MD,	DZ, IS, MG,	EC, JP, MK,	EE, KE, MN,	EG, KG, MW,	ES, KP, MX,	FI, KR, MZ,	GB, KZ, NA,	GD, LC, NI,	
	RW:	SY, BW, AZ, EE, RO,	TJ, GH, BY, ES, SE,	TM, GM, KG, FI, SI,	TN, KE, KZ, FR,	TR, LS, MD, GB, TR,	TT, MW, RU, GR, BF,	TZ, MZ, TJ, HU,	UA, NA, TM, IE,	UG, SD, AT, IS,	US, SL, BE, IT,	UZ, SZ, BG, LT,	VC, TZ, CH, LU,	VN, UG, CY, MC,	YU, ZM, CZ, NL,	ZA, ZW, DE, PL,	ZM, AM, DK, PT,	ZW
EP	1720 R:	AT, IS,	BE,	BG, LI,	CH, LT,	CY,	2006 CZ, MC,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	
PRIORIT	US 20070155754 A1 20070705 US 2006-598262 20060823 PRIORITY APPLN. INFO: W0 2005-047758P P 20040225 W0 2005-085491 W 20050222																	

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): CASREACT 143:266686; MARPAT 143:266686 GI

- AB Tetralins of formula I [R1 = CH2NR3R4, CONR3R4, NH-cycloalky1, N-methylpiperazinocarbonyl; R2 = H, NH-alky1, NR3R4, NH-cycloalky1, N-methylpiperazino, piperidino, pyrrolidino, etc.; R3 = H, alky1; R4 = alky1, phenylalkylene; R3R4 = alkylene, etc.] are prepared which have histamine-H3 receptor antagonist activity. The invention discloses pharmaceutical compns. comprising compds. of formula I as well as methods of using them to treat obesity and other histamine H3 receptor-related diseases. Thus, II was prepared and had Ki value of 1.5 nM against GTP y 135S1.
- IT 863925-32-0P 863925-33-1P R1: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RRCT (Reactant or reagent); USES (Uses) (preparation of tetralin derivs. as histamine H3 receptor antagonists) RN 863925-32-0 CAPJUS
- RN 863925-32-0 CAPLOS
 CN Methanone, [(2S)-2-(1-pyrrolidinylmethyl)-1-pyrrolidinyl](5,6,7,8-tetrahydro-2-naphthalenyl)- (CA INDEX NAME)

Absolute stereochemistry.

- RN 863925-33-1 CAPLUS
- CN Methanone, [(2R)-2-(1-pyrrolidinylmethyl)-1-pyrrolidinyl](5,6,7,8tetrahydro-2-naphthalenyl)- (CA INDEX NAME)

Absolute stereochemistry.

- IT 863925-34-2P 863925-35-3P RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
 - (Uses)
 (preparation of tetralin derivs. as histamine H3 receptor antagonists)
- RN 863925-34-2 CAPLUS
- CN Pyrrolidine, 2-(1-pyrrolidinylmethyl)-1-[(5,6,7,8-tetrahydro-2-naphthalenyl)methyl]-, (2S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 863925-35-3 CAPLUS

CN Pyrrolidine, 2-(1-pyrrolidinylmethyl)-1-[(5,6,7,8-tetrahydro-2-naphthalenyl)methyl]-, (2R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 3 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2004:354923 CAPLUS

DOCUMENT NUMBER: 140:375196

TITLE: Preparation of substituted piperazines,

[1,4]diazepines, and 2,5-diazabicyclo[2.2.1]heptanes as histamine H1 and/or H3 antagonists or histamine H3

reverse antagonists

Ancliff, Rachael; Eldred, Colin David; Fogden, Yvonne INVENTOR(S): C.; Hancock, Ashley Paul; Heightman, Thomas Daniel; Hobbs, Heather; Hodgson, Simon Teanby; Lindon, Matthew

J.; Wilson, David Matthew PATENT ASSIGNEE(S): Glaxo Group Limited, UK

SOURCE: PCT Int. Appl., 140 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

PA	PATENT NO.					KIND DATE				APP	LICAT	ION I	NO.		D	ATE	
WO	2004	0355	56		A1		2004	0429		WO.	2003-	EP11	423		2	0031	014
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		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	, EE,	EG,	ES,	FI,	GB,	GD,	GE,
		GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JΡ	, KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,
											, MN,						
											, SE,						TM,
											, VN,						
	RW:										, TZ,						
											, СН,						
											, NL,						
	0500										, GW,						
CA	2502	249	0.0		AI		2004	0429		CA	2003- 2003-	2502.	249		2	0031	014
AU	2003	2803	80		AI		2004	0504		AU	2003-	2803	80		2	0031	014
BR	2003	0152	83		A		2005	0830		BR	2003- 2003-	1528.	3		2	0031	014
EP	120/	211	DE	OII	AI	DIZ	2005	0831	CD	EP	, IT,	1122.	Z I	NIT	CE Z	0031	UI4
CN	1726	201	51,	ш.,	ъ,	, FI, RO, MK, 20060125 20080709 20060316 20061130 20071114 20080328 2 20080710 20060224 20050707				CM MΠ	2003-	8010	6014	EE,	2	0031	014
CM	1004	0052	3		Ċ		2008	0709		CIT	2005	0010	0011		-	0031	014
JP	2006	5089	3.5		T		2006	0316		JP	2004-	5442	41		2	0031	014
NZ	5394	46			Ā		2006	1130		NZ	2003-	5394	46		2	0031	014
CN	1010	7030	9		A		2007	1114		CN	2006-	1010	8610		2	0031	014
NZ	5499	63			A		2008	0328		NZ	2003-	5499	63		2	0031	014
RU	2328	494			C2		2008	0710		RU	2005-	1100	61		2	0031	014
IN	2005	KN00	566		A		2006	0224		IN	2005-1	KN56	6		2	0050	404
ZA	2005	0028	73		A		2006	0726		ZA	2005-	2873			2	0050	408
US	2006	0025	404		A1		2006	0202		US	2005-	5317.	58		2	0050	414
US	7615	550			B2		2009	1110									
MX	2005	0040	78		A		2005	0608		MX	2005-	4078			2	0050	415
ZA	2006	0036	04		A		2007	0425		ZA	2006-	3604			2	0060	505
IN	2006	KN02	281		A		2007	0525		TM	2006-1	KN22:	RT		2	0060	RIO
US US MX ZA IN JP PRIORIT	∠007	0160	41		A		2007	0125		O.D.	2006-	2311	0.3		. 2	0060	828
PRIORIT	1 APP	TM.	TMEO	. :						GB	2002-: 2003-:	2408	4 CO 1 4		A 2	0021	014
											2003-						
										UP	2004-	J44Z	4.1		ms Z	0031	0.1.4

NZ 2003-539446 A3 20031014 WO 2003-EP11423 W 20031014 IN 2005-KN566 A3 20050404

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 140:375196

$$\begin{bmatrix} R^1 \\ Z \\ N \end{bmatrix}_p \begin{bmatrix} R^2 \\ N \end{bmatrix}_n \begin{bmatrix} R^14 \\ K \end{bmatrix}_k \begin{bmatrix}$$

AB The title compds. [I; R1 = H, alkvl, alkoxv, etc.; Z = a bond, CO, (un) substituted CONH, SO2; p = 1-2; m, n, r = 0-2; R2 = halo, alkvl, alkoxy, etc.; R3 = (CH2)gNR11R12, II (wherein g = 2-4; R11, R12 = alkyl, cycloalkyl; NR11R12 = heterocyclyl; R13 = H, alkyl, cycloalkyl, etc.; R14 = halo, alkyl, haloalkyl, etc.; f, k = 0-2; q = 0-2; h = 0-3, such that qand h cannot both be 0); R4 = H, alkyl such that when r = 2, two R4 groups may instead be linked to form CH2, (CH2)2, (CH2)3; with the provisos], useful in the treatment of neurodegenerative disorders including Alzheimer's disease, and inflammatory diseases of the upper respiratory tract, were prepared Thus, reacting 1-[4-(3-piperidin-1-ylpropoxy)benzyl]piperazine.3HCl (preparation given) with benzoic acid afforded 77% III which was tested in the histamine H3 functional antagonist assay and showed pKb of > 6.5. The pharmaceutical composition comprising the compound I is claimed. 684244-55-1P 684244-76-6P 684244-95-9P 684245-53-2P 684245-17-8P 684245-35-0P

084249-17-8F 084249-33-UF 084249-33-4F 084249-33-4F 084249-33-4F 084249-31-4F 08424

(preparation of substituted piperazines, [1,4]diazepines, and

2,5-diazabicyclo[2.2.1]heptanes as histamine H1 and/or H3 antagonists or histamine H3 reverse antagonists)

RN 684244-55-1 CAPLUS

CN Methanone, [4-[4-[3-(1-piperidiny1)propoxy]pheny1]-1-piperaziny1](5,6,7,8-tetrahydro-2-naphthaleny1)- (CA INDEX NAME)

RN 684244-76-6 CAPLUS

CN Methanone, [4-[4-[3-(5-ethyl-2-methyl-1-piperidinyl)propoxy]phenyl]-1-piperazinyl](5,6,7,8-tetrahydro-2-naphthalenyl)- (CA INDEX NAME)

RN 684244-95-9 CAPLUS

CN Methanone, [4-[4-[3-(2-methyl-1-piperidinyl)propoxy]phenyl]-1-piperazinyl](5,6,7,8-tetrahydro-2-naphthalenyl)-, 2,2,2-trifluoroacetate (1:1) (CA INDEX NAME)

CM :

CRN 684244-94-8 CMF C30 H41 N3 O2

10/513699

CM 2

CRN 76-05-1 CMF C2 H F3 O2

F-C-CO₂H

RN 684245-17-8 CAPLUS

CN Methanone, [4-[4-[3-(hexahydro-1(2H)-azociny1)propoxy]phenyl]-1-piperazinyl](5,6,7,8-tetrahydro-2-naphthalenyl)-, 2,2,2-trifluoroacetate (1:1) (CA INDEX NAME)

CM 1

CRN 684245-16-7

CMF C31 H43 N3 O2

CM 2

CRN 76-05-1 CMF C2 H F3 O2

$${\scriptstyle F-C-CO_2H\atop \downarrow}$$

CN

RN 684245-35-0 CAPLUS

Methanone, [4-[4-[3-(cyclopentylmethylamino)propoxy]phenyl]-1piperazinyl](5,6,7,8-tetrahydro-2-naphthalenyl)- (CA INDEX NAME)

Erich Leese

<12/04/2007>

RN 684245-53-2 CAPLUS

CN Methanone, [4-[4-[3-(3-methyl-1-piperidinyl)propoxy]phenyl]-1-piperazinyl](5,6,7,8-tetrahydro-2-naphthalenyl)- (CA INDEX NAME)

OS.CITING REF COUNT:

14 THERE ARE 14 CAPLUS RECORDS THAT CITE THIS

RECORD (19 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 4 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2004:333695 CAPLUS

DOCUMENT NUMBER: 140:339199

TITLE: Preparation of 1,4-disubstituted piperidine

derivatives and their use as 11-βHSD1 inhibitors INVENTOR(S): Barton, Peter John; Jewsbury, Philip John; Pease,

Janet Elizabeth

PATENT ASSIGNEE(S): Astrazeneca Ab, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 144 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.									ICAT				D.	ATE			
WO											2003-				2	0031	007
	W:										BG,						
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	GE,
		GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,
		LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NI,	NO,	ΝZ,
		OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ΤJ,	TM,
											VN,						
	RW:										TZ,						
											CH,						
											NL,						
											GW,						
		2501611 A1 20040422 2003269242 A1 2004050															
EP											2003-						
	R:										IT,						
											TR,						
BR	2003	0151	66		A		2005	0816		BR 2	2003-	1516	6		2	0031	007
CN	1723	199			A		2006	0118		CN 2	2003-	8010	5353		2	0031	007
JP	2006	5064	51		Т		2006	0223		JP 2	2005-	5009	93		2	0031	007
	2005																
	2005																
					A		2006	0222									
KIT!	APP	LN.	TNEO	. :							2002-						
											2003-					0030	
										WO 2	2003-0	GB43	TR	1	71 2	0031	007

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 140:339199

GI

$$\begin{bmatrix} R^{12} \end{bmatrix}_{n} \\ Q \\ \begin{bmatrix} R^{12} \end{bmatrix}_{m} \\ N \\ X \\ \end{bmatrix}$$

DM

- AB The title compds. II; A = carbocyclyl, heterocyclyl; Rl = halo, NO2, CN, OH, etc.; n = 0-5; X = a bond, CO, SO2, CONRIL, CSNRI, C(0)O, C(NRII), CH2 (wherein Rl1 = H, alkyl, carbocyclyl, heterocyclyl); Y = H, alkyl, alkenyl, carbocyclyl, etc.; Rl2 = OH, Me, Et. Pr; m, q = 0-1], useful in the manufacture of a medicament for treating diabetes, obesity, hyperlipidemia, etc., were prepared Thus, reacting (4-chlorophenyl)(4-piperidyl)methanone. HCl with 4-fluorobenzoyl)-chloride in the presence of Et3N in DCM afforded 29% 1-(4-fluorobenzoyl)-4-(4-chlorobenzoyl)piperidine. The compds. I typically show an IC50 < 10 μM against 11βHSDl. The pharmaceutical composition comprising the compound I is claimed.
- IT 681130-55-2P Rl: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of 1,4-disubstituted piperidine derivs. and their use as $11-\beta HSD1$ inhibitors)

- 681130-55-2 CAPLUS
- CN Piperidine, 4-(4-fluorobenzoy1)-1-[(5,6,7,8-tetrahydro-1-naphthaleny1)carbony1]- (9CI) (CA INDEX NAME)

- OS.CITING REF COUNT: 19 THERE ARE 19 CAPLUS RECORDS THAT CITE THIS RECORD (29 CITINGS)
- REFERENCE COUNT: 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2003:696782 CAPLUS

DOCUMENT NUMBER: 139:230625

TITLE: Preparation of bipiperidinyl and related compounds as acetyl CoA carboxylase inhibitors useful against

metabolic syndrome and other disorders

INVENTOR(S): Perry, David Austen; Harwood, Harold James, Jr.

PATENT ASSIGNEE(S): Pfizer Products Inc., USA

SOURCE: PCT Int. Appl., 181 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

P.	PATENT NO.				KIND DATE						LICAT					ATE	
W	0 200	30721	97													0030	217
	W:										, BG,						
											, EE,						
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE	, KG,	KP,	KR,	KZ,	LC,	LK,	LR,
											, MW,						
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SK,	SL	, TJ,	TM,	TN,	TR,	TT,	TZ,	UA,
		UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	zw								
	RW										, TZ,						
											, CH,						
											, NL,						BF,
											, ML,						
	U 200																
	P 147									EP :	2003-	7428	82		2	0030	217
E	P 147																
	R:										, IT,						PT,
											, TR,						
C	N 164	2599			A		2005	0720		CN :	2003-	8069	90		2	0030	217
A	T 303	178			T		2005	0915		AT :	2003-	7428	82		2	0030	217
	S 224																
	Z 534.																
	S 200									US :	2003-	3708	44		2	0030	220
	S 697				B2		2005										
	N 200																
	A 200																
					A		2004	1124	24 NO 2004-4034 US 2002-365358P								
PRIORI	TY API	PLN.	INFO	.:													
										WO :	2003-	IB57	3	1	W 2	0030:	217

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 139:230625 GI

Acetyl CoA carboxylase (ACC) inhibitors (shown as I; variables defined below; most examples include the bipiperidinyl ring system, e.g. (anthracen-9-yl)[(3R)-3-(morpholine-4-carbonyl)[1,4']bipiperidinyl-1'yl]methanone), pharmaceutical compns. containing such compds. and the use of such compds. to treat for example, Metabolic Syndrome, atherosclerosis, diabetes and obesity are disclosed. None of pharmacol. activity, therapeutic uses and methods of preparation is claimed and pharmacol. data are not included. More than 200 example prepns. and/or characterization data are included for I and intermediates. For I: A-B is N-CH or CH-N; K is (CH2)r (r = 2-4); m and n = 1-3 when A-B is N-CH or 2 or 3 when A-B is CH-N; the dashed line = the presence of an optional double bond; D is carbonyl or sulfonyl. E is either (a) a bicyclic ring consisting of two fused fully unsatd. 5-7 membered rings, taken independently, each of said rings optionally having 1-4 heteroatoms = O, S and N; or (b) a tricyclic ring consisting of two fused fully unsatd. 5-7 membered rings, taken independently, each of said rings optionally having 1-4 heteroatoms = 0, S and N, said two fused rings fused to a 3rd partially saturated, fully unsatd. or fully saturated 5-7 membered ring, said 3rd ring optionally having 1-4 heteroatoms = O, S and N. Or (c) a tetracyclic ring comprising a bicyclic ring consisting of two fused fully unsatd. 5-7 membered rings, taken independently, each of said rings optionally having 1-4 heteroatoms = 0, S and N, said bicyclic ring fused to two fully saturated, partially saturated or fully unsatd. 5-7 membered monocyclic rings taken independently, each of said rings optionally having 1-4 heteroatoms = O, S and N or said bicyclic ring fused to a 2nd bicyclic ring consisting of two fused fully saturated, partially saturated or fully unsatd. 5-7 membered rings, taken independently, each of said rings optionally having 1-4 heteroatoms = 0, S and N; or (d) a teraryl ring comprising a fully unsatd. 5-7 membered ring, said ring optionally having 1-4 heteroatoms = O, S and N, and said ring disubstituted independently with a fully unsatd. 5-7 membered ring to form a teraryl nonfused ring system, each of said substituent rings optionally having 1-4 heteroatoms = O, S and N. G is carbonvl, sulfonvl or CR7R8 (R7 and R8 = H, (C1-C6)alkyl, (C2-C6) alkenyl or (C2-C6)alkynyl or a 5-7 membered partially saturated, fully saturated or fully unsatd. ring optionally having one heteroatom = O, S and N); J is OR1, NR2R3 or CR4R5R6; addnl. details including provisos are given in the claims. 591781-07-6P, 1'-(1,2,3,4-Tetrahydroanthracen-9-

<12/04/2007> Erich Leese

ylcarbonyl)[1,4']bipiperidinyl-3-carboxylic acid diethylamide

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug candidate; preparation of bipiperidinyl and related compds. as acetyl CoA carboxylase inhibitors useful against metabolic syndrome and other disorders)

RN 591781-07-6 CAPLUS

CN [1,4'-Bipiperidine]-3-carboxamide,

N,N-diethyl-1'-[(1,2,3,4-tetrahydro-9-anthracenyl)carbonyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 16 THERE ARE 16 CAPLUS RECORDS THAT CITE THIS RECORD (30 CITINGS)

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

I.4 ANSWER 6 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2000:91070 CAPLUS

DOCUMENT NUMBER: 132 - 166198

TITLE: Synthesis and platelet aggregation inhibitory activity of 6- [(4-substituted-piperazinyl)phenyl]-5-methyl-4,5-

dihydro-3(2H)pyridazinones

Wu, Qiuye; Ni, Jin; Jiang, Yuanying; Liu, Chaomei; Wu, AUTHOR(S):

Bo; Zhang, Guangming; Yao, Jiayong

CORPORATE SOURCE: Faculty of Pharmacy, Second Military Medical University, Shanghai, 200433, Peop. Rep. China

SOURCE: Zhongguo Yaowu Huaxue Zazhi (1999), 9(4), 259-263 CODEN: ZYHZEF; ISSN: 1005-0108

PUBLISHER: Zhongguo Yaowu Huaxue Zazhi Bianjibu

DOCUMENT TYPE: Journal

LANGUAGE: Chinese

CH 3

Title compds. I (R = CH3, CH3CH2, CH3(CH2)3, (CH3)2CHCH2CH2, CH3(CH2)7, AB CH3(CH2)11, CH3(CH2)15, C6H5CH2, 4-C1C6H4CH2, 2-C1C6H4CH2, 3-C1C6H4CH2, 4-CH3C6H4CH2, CH3OCOCH2, 4-CH3CH2OCO-C6H4CH2) were prepared from N-acetylaniline via acylation, hydrolysis, cyclization and substitution. The results of preliminary pharmacol. tests showed that all the synthetic compds. had activity against platelet aggregation induced by ADP in vitro in rabbits.

259140-66-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(synthesis and platelet aggregation inhibitory activity of

6-methyl-6-piperazinylphenyldihydropyridazinones)

RN 259140-66-4 CAPLUS

CN 3(2H)-Pyridazinone, 4,5-dihydro-5-methyl-6-[4-[4-[4-[(5,6,7,8-tetrahydro-2naphthalenyl)methyl]-1-piperazinyl]phenyl]- (CA INDEX NAME)

Me CH2-

L4 ANSWER 7 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 2000:62206 CAPLUS

DOCUMENT NUMBER: 132:207835

TITLE: Regioselective aminomethylations of bicyclic phenols AUTHOR(S): Lange, Jos; Hoogeveen, Sonja; Veerman, Willem; Wals,

Henri

CORPORATE SOURCE: Medicinal Chemistry Department, Solvay Pharmaceuticals
Research Laboratories, Weesp, 1380 DA, Neth.

SOURCE: Research Laboratories, weesp, 1380 1 800 SOURCE: Heterocycles (2000), 53(1), 197-204

CODEN: HTCYAM; ISSN: 0385-5414

PUBLISHER: Japan Institute of Heterocyclic Chemistry

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 132:207835

AB The regioselectivity in the aminomethylation, Mannich reaction, of bicyclic phenols was studied. Highly regioselective Mannich reactions

enable easy synthetic access to novel bicyclic [(dialkylamino)methyl]phenols under very mild reaction conditions.

IT 260394-47-6P 260394-48-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(regioselective aminomethylation of bicyclic phenols)

RN 260394-47-6 CAPLUS

CN 2-Naphthalenol, 5,6,7,8-tetrahydro-3-[(4-methyl-1-piperazinyl)methyl]-(CA INDEX NAME)

RN 260394-48-7 CAPLUS

CN 2-Naphthalenol, 5,6,7,8-tetrahydro-1-[(4-methyl-1-piperazinyl)methyl]-(CA INDEX NAME)

OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

REFERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 8 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1999:729780 CAPLUS

DOCUMENT NUMBER: 132:222471

TITLE: Synthesis and platelet aggregation activity of 6-[4-substituted-piperazinyl)phenyl]-4,5-dihydro-3(2H)-

AUTHOR(S): Wu, Oiuve; Zhang, Guangming; Liao, Hongli; Liu,

AUTHOR(S): Wu, Qiuye; Zhang, Guangming; Liao, Hongli; Liu,
Chaomei

CORPORATE SOURCE: Faculty of Pharmacy, Second Military Medical Univ., Shanghai, 200433, Peop. Rep. China

SOURCE: Zhongguo Yaowu Huaxue Zazhi (1999), 9(3), 172-175, 185

CODEN: ZYHZEF; ISSN: 1005-0108
PUBLISHER: Zhongquo Yaowu Huaxue Zazhi Bianjibu

PUBLISHER: Zhongguo Yaowu Huaxue Zazhi Bianjibu

LANGUAGE: Southai

CT

$$R-N \longrightarrow N-N \longrightarrow N-N \longrightarrow N-N$$

AB Bighteen title compds. I (R = CH3, CH3CH2, CH3(CH2)3, (CH3)2CHCH2CH2, CH3(CH2)7, CH3(CH2)11, CH3(CH2)12, CH3(CH3)12, CH3(CH

260979-38-2P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(synthesis and platelet aggregation activity of substituted piperazinylphenyldihydropyridazinones)

RN 260979-38-2 CAPLUS

CN 3(2H)-Pyridazinone, 4,5-dihydro-6-[4-[4-[4-[(5,6,7,8-tetrahydro-2-naphthalenyl)methyl]-1-piperazinyl]phenyl]- (CA INDEX NAME)

L4 ANSWER 9 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1999:511159 CAPLUS

DOCUMENT NUMBER: 131:157709

TITLE: Preparation of bicyclic pyridine and pyrimidine derivatives as neuropeptide Y receptor antagonists

INVENTOR(S): Norman, Mark H.; Chen, Ning; Han, Nianhe; Liu, Longbin; Hurt, Clarence R.; Fotsch, Christopher H.;

Jenkins, Tracy J.; Moreno, Ofir A.

PATENT ASSIGNEE(S): Amgen Inc., USA

SOURCE: PCT Int. Appl., 469 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English

LANGUAGE: Engl FAMILY ACC. NUM. COUNT: 1

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.															ATE		
	9940															 9990	205
	W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR	, BY,	CA,	CH,	CN,	CU,	CZ,	DE,
		DK,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM	, HR,	HU,	ID,	IL,	IN,	IS,	JP,
		KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS	, LT,	LU,	LV,	MD,	MG,	MK,	MN,
		MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD	, SE,	SG,	SI,	SK,	SL,	TJ,	TM,
		TR,	TT,	UA,	UG,	UZ,	VN,	YU,	ZW								
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW	, AT,	BE,	CH,	CY,	DE,	DK,	ES,
		FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL	, PT,	SE,	BF,	BJ,	CF,	CG,	CI,
		CM,	GA,	GN,	GW,						, TG						
US	6187	777			B1		2001	0213		US	1999-	2467	75		1	9990	204
CA	2319	275			A1		1999	0812		CA	1999-	2319	275		1	9990	205
					C 20071016												
AU	9926	590				A 19990823 AU 1999-26590									1	9990	205
	7479				B2 20020530												
EP	1054	887								ΕP	1999-	9067	56		1	9990	205
EP	1054				B1		2006										
	R:								GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,
		ΙE,	SI,	LT,	LV,	FI,	RO,										
JP	2003 3230	5022	72		T						2000-					9990	
AT	3230	88			T			0415			1999-					9990	
ES	2257	851			Т3		2006				1999-					9990	
	9900							0806		ZA	1999-	967			1	9990	
	2000							0219			2000-					0000	
	6583				B1		2003	0624			2000-					0000	
PRIORIT	Y APP	LN.	INFO	.:							1998-						
											1998-					9980	
											1998-					9980	
									US 1998-93577P						9980		
											1999-					9990	
									WO 1999-US2500				00		W 1	9990	205

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 131:157709

GI

Ι

AB Title compds.[I; R = H, CH3, (CH3)2CH, SCH3, CH3CH2, NH2, CF3, NHCOC6H5, cyclopropyl, CH2OH, (CH3)2CH2CH2, N(CH3)2, OCH3, NHCH3, NH(CH2)4NH2; R1 = NH, S, NCH3, O; R2 = H, COCH3, C6H5, CH3, CH3CH2; R3 = NH2, CH3, NHC6H5, N(CH2CH3)2, (CH3CH2)N(CH2)3CH3, (CH3)N(CH2)2NHCH3, N(CH3)CH(CH3)CH(Ph)OH, (CH3CH2)NCH2C(CH3):CH2, NHCH2CF3, NHCH2CH2C6H5, NH(CH2)3OCH2CH3, 4-ClC6H4, 4-CH3OC6H5, 2-thienyl, 1-pyrrolidinyl, 1-piperidinyl, 4-morpholinyl, 1-piperazinyl, 3-pyridyl; R4 = C6H5, 4-CH3C6H4, 4-ClC6H4, (CH3)3C, 4-FC6H4, 3-HOC6H4, 2-pyridyl, cyclohexyl, 2-furyl, 2-FC6H4 2-thienyl, 1-adamantyl, CH3, 4-CH3OC6H4; X = N, CH; etc.], pharmaceutical acceptable salts, ester, solvate, and N-oxide are prepared and tested as neuropeptide Y receptor antagonists in the modulation of feeding behavior, obesity, diabetes, cancer, inflammatory disorders, depression, stress related disorders, Alzheimer's disease and other disease conditions. Thus, the title compound I (R = CH3; R1 = NH; X = N; R2 = H; R3 = N(CH2CH3)2; R4 = C6H5) was prepared

237436-39-4

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of pyrrolopyridine and pyrrolopyrimidine derivs. as neuropeptide Y receptor antagonists)

RN 237436-39-4 CAPLUS

CN Pyrrolidine, 1-[1-(5,6,7,8-tetrahydro-2-naphthalenyl)ethenyl]- (CA INDEX NAME)

OS.CITING REF COUNT:

THERE ARE 25 CAPLUS RECORDS THAT CITE THIS RECORD (29 CITINGS)

- REFERENCE COUNT:
- THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

<12/04/2007> Erich Leese

25

L4 ANSWER 10 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1999:495272 CAPLUS

DOCUMENT NUMBER: 131:130011

TITLE: Preparation of N-acyl-2-aminoacetamides and

cyclization products thereof.

INVENTOR(S): Hulme, Christopher; Morton, George C.; Salvino, Joseph

APPLICATION NO.

DATE

M.; Labaudiniere, Richard F.; Mason, Helen J.;
Morrissette, Matthew M.; Ma, Liang; Cherrier,

Marie-Pierre

PATENT ASSIGNEE(S): Rhone-Poulenc Rorer Pharmaceuticals Inc., USA

KIND DATE

SOURCE: PCT Int. Appl., 156 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PAIENI NO. KI										APP	PICMI		D	MIE			
WO	9938	844			A1		1999	0805		WO	1999-	US19	23		1	9990	129
	W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR	, BY,	CA,	CN,	CU,	CZ,	DE,	DF
											, JP,						
		LK,	LR,	LS,	LT,	LU,	LV,	MD,	MG,	MK	, MN,	MW,	MX,	NO,	NZ,	PL,	PI
		RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ	, TM,	TR,	TT,	UA,	UG,	US,	U2
			YU,														
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	UG,	ZW	, AT,	BE,	CH,	CY,	DE,	DK,	E:
		FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL	, PT,	SE,	BF,	ВJ,	CF,	CG,	C:
		CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD	, TG						
CA	2318	601			A1		1999	0805		CA	1999- 1999-	2318	601		1	9990	129
ΑU	9924	821			A		1999	0816		AU	1999-	2482	1		1	9990	129
AU	7479	87			B2		2002	0530									
za	9900	729			A		2000	0110		ZA	1999-	729			1	9990	12
EP	1051	397			A1		2000	1115		EP	1999-	9044	21		1	9990	12
EΡ	1051	397			B1		2008	1231									
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	Ρ
		IE,	SI,	FI,	RO,	CY											
BR	9908	207			A		2000	1128		BR	1999- 2000- 2001- 1999- 2000-	8207			1	9990	12
JΡ	2002	5019	44		T		2002	0122		JΡ	2000-	5300	81		1	9990	12
HU	2001	0013	29		A2		2002	0328		HU	2001-	1329			1	9990	12
HU	2001	0013	29		A3		2002	0729									
CN	1173	946			C		2004	1103		CN	1999-	8025	03		1	9990	12
ΑP	1462				A		2005	0930		AP	2000-	1864			1	9990	12
IL	1375	71			A		2006	1210		IL	1999-	1375	71		1	9990	12
ΑT	4192	:33			T		2009	0115		ΑT	1999-	9044	21		1	9990	12
US	6492	553			B1		2002	1210		US	1999- 1999- 1999-	3682	13		1	9990	80
NO	2000	0037	92		A		2000 2007	0927		NO	2000-	3792			2	0000	72
NO	3240	67			B1		2007	0806									
MX	2000	0075	55		A		2001	0219			2000-						
BG	1047	24			A		2001	0330		BG	2000-	1047	24		2	0000	82
BG	6505	7			B1		2007	0131									
RIT:	Y APP	LN.	INFO	.:			2007				1998-						
											1998-						
										US	1998-	9870	8P		A2 1	9980	90
										US	1998- 1999-	1010	56P		A2 1	9980	91
										WO	1999-	US19	23		W 1	9990	129

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OTHER SOURCE(S): MARPAT 131:130011

AB RaRbNCRcaRcbRd Ra = RaaCO; Dd = CONHRda; Raa, Rb, Rca, Rcb = H, (substituted) aliphatyl, aryl; Rda = (substituted) aliphatyl, aryl; with provisos were prepared by reaction of RcaCORcb with RbNH2, RaCO2H, and NCRda. Title compds. may be prepared on a isocyanide resin and deprotected/cyclized to give 1,4-benzodiazepine-2,5-diones, diketopiperazines, ketopiperazines, lactams, 1,4-benzodiazapines, and dihydroquinoxalinones.

IT 234781-39-6P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of N-acyl-2-aminoacetamides and cyclization products thereof)
RN 234*RB-139-6 CAPLUS

CN 2-Piperazinepentanoic acid, 3-oxo-1-[(5,6,7,8-tetrahydro-2-naphthalenyl)carbonyl]-, methyl ester (CA INDEX NAME)

OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (5 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 11 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1999:172595 CAPLUS

DOCUMENT NUMBER: 130:223167

TITLE: Preparation of piperidinylpyrrolidins as modulators of

chemokine receptor activity

INVENTOR(S): Budhu, Richard J.; Hale, Jeffrey J.; Holson, Edward; Lynch, Christopher; Maccoss, Malcolm; Mills, Sander

G.; Berk, Scott C.; Willoughby, Christopher A.

PATENT ASSIGNEE(S): Merck & Co., Inc., USA PCT Int. Appl., 262 pp.

SOURCE:

CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.					KIND DATE					ICAT:	ION	NO.		D,	ATE	
WO 99	909984			A1	-	 1999	0304		WO 1	998-	US17	755		1	9980	827
Ţ.	V: AL,	AM,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CN,	CU,	CZ,	EE,	GE,	HR,
	HU,	ID,	IL,	IS,	JP,	KG,	KR,	KZ,	LC,	LK,	LR,	LT,	LV,	MD,	MG,	MK,
	MN,	MX,	NO,	NZ,	PL,	RO,	RU,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	UA,
	US,	UZ.	VN.	YU												
F	RW: GH,	GM.	KE.	LS.	MW.	SD.	SZ.	UG.	ZW.	AT.	BE.	CH.	CY.	DE,	DK.	ES.
	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	BJ,	CF,	CG,	CI,
	CM,	GA,	GN,	GW,	ML,	MR.	NE.	SN,	TD,	TG						
CA 22	CA 2298813					1999	0304		CA 1	998-	2298	813		1	9980	827
AU 98	392067			A		1999	0316		AU 1	998-	9206	7		1	9980	827
EP 10	09405			A1		2000	0621		EP 1	998-	9445	48		1	9980	827
F	R: AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	PT,	IE,
	SI,	LT,	LV,	FI,	RO											
US 61	166037			A		2000	1226		US 1	998-	1412	27		1	9980	827
JP 20	015261	78		T		2001	1218		JP 2	000-	5073	74		1	9980	827
PRIORITY A	APPLN.	INFO	. :						US 1	997-	5774	3P	I	2 1	9970	828
									GB 1	998-	1009		2	A 1	9980	116
						WO 1998-US17755						1 1	9980	827		

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 130:223167 GI

$$R^2 = -N$$

$$R^2 = -N$$

$$0 = N$$

$$1 = N$$

AB Title modulators [I; Rl = CH2Ph, SO2Ph, CONHPh, H, COPh, (CH2)3Ph,
l-fluorenecarbonyl, etc.; R = OH, H, Ph, CF3, CH2Ph, etc.; n = 0-2; S = S,
C; R2 = benzo[d]azepin-3-yl, 4-phenyl-perhydroazepin-1-yl, etc.],
pharmaceutically acceptable salts thereof, individual diastereomers, and
enantiomers thereof are prepared as modulators of chemokine receptor
activity. 21X19 combinatorial library was mentioned using com. available
4-sulfamylbenzoyl polystyrene resin supported subunits (21 pools) of
trifluoromethylsulfonyl chloride, arylsulfonyl(carbonyl) chlorides, and
heterocyclic sulfonyl(carbonyl) chlorides. Thus, compound II was prepared
from Me (Z)-cinnamate and N-(methoxymethyl)-N-
(trimethylsilvlumethyl)benzylamine via seven steps.

T 221141-11-3P 221157-12-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation of piperidinylpyrrolidins as modulators of chemokine receptor activity)

- RN 221141-11-3 CAPLUS
- CN Methanone, [(3R,4S)-3-phenyl-4-[(4-phenyl-1-piperidinyl)methyl]-1pyrrolidinyl](5,6,7,8-tetrahydro-1-naphthalenyl)-, rel- (CA INDEX NAME)

Relative stereochemistry.

RN 221157-12-6 CAPLUS

CN Methanone, [(3R,4R)-3-phenyl-4-[(4-phenyl-1-piperidinyl)methyl]-1pyrrolidinyl](5,6,7,8-tetrahydro-1-naphthalenyl)-, rel- (CA INDEX NAME)

Relative stereochemistry.

OS.CITING REF COUNT: 34 THERE ARE 34 CAPLUS RECORDS THAT CITE THIS RECORD (34 CITINGS)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 12 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1998:220858 CAPLUS DOCUMENT NUMBER: 128:270614

ORIGINAL REFERENCE NO.: 128:53569a,53572a

TITLE: Preparation of acylpiperazines and related compounds

as inhibitors of farnesyl-protein transferase.

INVENTOR(S): Graham, Samuel L.; Williams, Theresa M. PATENT ASSIGNEE(S): Merck and Co., Inc., USA

SOURCE: U.S., 50 pp., Cont.-in-part of U.S. Ser. No. 237,586,

abandoned. CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT	KIND DATE					APPL	ICAT	ION	NO.		D.	ATE			
		-		_									-		
US 5736	539		A		1998	0407		US 1	995-	5498	29		1	9951	116
WO 9500	497		A1		1995	0105		WO 1	994-	US56	34		1	9940	519
W:	AU, BB,	BG,	BR,	BY,	CA,	CN,	CZ,	FI,	GE,	HU,	JP,	KG,	KR,	KZ,	LK,
	LV, MD, MG,				NO,	NZ,	PL,	RO,	RU,	SD,	SI,	SK,	TJ,	TT,	UA,
	US, UZ														
RW:	AT, BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,
	BF, BJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG		
ZA 9404	326		A		1995	1214		ZA 1	994-	4326			1	9940	617
PRIORITY APP).:						US 1	993-	8002	8		B2 1	9930	618	
							US 1994-237586						B2 1	9940	511
									WO 1994-US5634						519

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 128:270614 GT

Ι

- AB Title compds. e.g., [I; X = 0, H2; m = 1, 2; n = 0, 1; t = 1, 4; R, R1 = H, alkyl, aralkyl; R2-R5 = H, (substituted) alkyl, alkenyl, alkynyl, aryl, heterocyclyl, acyl; Y = (substituted) aryl, heterocyclyl], were prepared Thus, 1-[2(R)-amino-3-mercaptopropy1]-2(S)-[2-(3-pyridylmethoxy)ethy1]-4-(1-naphthoyl)piperazine trihydrochloride (preparation given) inhibited RAS farnesylation with IC50 = 1 nM.
- 169449-54-1 1099473-75-2 RL: PRPH (Prophetic)

(Preparation of acylpiperazines and related compounds as inhibitors of farnesvl-protein transferase.)

- RN 169449-54-1 CAPLUS
- 1-Piperazinepropanethiol, β-amino-2-(2-methoxyethyl)-4-[(5,6,7,8tetrahydro-1-naphthalenyl)carbonyl]-, [S-(R*,S*)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 1099473-75-2 CAPLUS CN INDEX NAME NOT YET ASSIGNED

Absolute stereochemistry.

IT 169449-55-2P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of acylpiperazines and related compds. as inhibitors of

farnesyl-protein transferase)

RN 169449-55-2 CAPLUS CN 1-Piperazinepropane

1-Piperazinepropanethiol, β -amino-2-(2-methoxyethyl)-4-[(5,6,7,8-tetrahydro-1-naphthalenyl)carbonyl]-, [S-(R*,S*)]-, bis(trifluoroacetate) (salt) (9C1) (CA INDEX NAME)

CM 1

CRN 169449-54-1 CMF C21 H33 N3 O2 S

Absolute stereochemistry.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

OS.CITING REF COUNT:

5

REFERENCE COUNT:

THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (5 CITINGS)

24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 13 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1996:70133 CAPLUS DOCUMENT NUMBER: 124:164423

ORIGINAL REFERENCE NO.: 124:30167a,30170a

TITLE: Synthesis and antimalarial activity of Mannich bases of N-tetrahydronaphthol-substituted 9-amino acridines

AUTHOR(S): Cao, Shouhai; Li, Fulin

CORPORATE SOURCE: Institute of Pharmacology and Toxicology, Academy of Military Medical Sciences, Beijing, 100071, Peop. Rep.

SOURCE: Zhongguo Yiyao Gongye Zazhi (1995), 26(7), 292-4 CODEN: ZYGZEA; ISSN: 1001-8255

PUBLISHER: Zhongguo Yivao Gongye Zazhi Bianjibu

DOCUMENT TYPE: Journal

LANGUAGE: Chinese

AB Nine Mannich bases of N-tetrahydronaphthol-substituted 9-amino acridines (I; R = Me, Et, Pr, iso-Pr, Bu, iso-Bu, sec-Bu, pentyl, isopentyl) were synthesized by using α-naphthol and 2-methoxy-6,9-dichloroacridine as starting materials. Preliminary screening showed that the suppressive activity of I (R = Bu, iso-Bu, sec-Bu) against P. berghei was equivalent to that of chloroquine and all the others were inferior to chloroquine.

173739-07-6P 173739-08-7P 173739-09-8P 173739-10-1P 173739-11-2P 173739-12-3P 173739-13-4P 173739-14-5P 173739-15-6P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(synthesis and antimalarial activity of Mannich bases of tetrahydronaphthol-substituted amino acridines)

Ι

173739-07-6 CAPLUS RN

CN 1-Naphthalenol, 4-[(6-chloro-2-methoxy-9-acridinyl)amino]-5,6,7,8tetrahydro-2-[(4-methyl-1-piperazinyl)methyl]- (CA INDEX NAME)

RN 173739-08-7 CAPLUS

CN 1-Naphthalenol, 4-[(6-chloro-2-methoxy-9-acridinyl)amino]-2-[(4-ethyl-1-piperazinyl)methyl]-5,6,7,8-tetrahydro- (CA INDEX NAME)

RN 173739-09-8 CAPLUS

CN 1-Naphthalenol, 4-[(6-chloro-2-methoxy-9-acridinyl)amino]-5,6,7,8tetrahydro-2-[(4-propyl-1-piperazinyl)methyl]- (CA INDEX NAME)

RN 173739-10-1 CAPLUS

CN 1-Naphthaleno1, 4-[(6-chloro-2-methoxy-9-acridiny1)amino]-5,6,7,8tetrahydro-2-[[4-(1-methylethy1)-1-piperaziny1]methy1]- (CA INDEX NAME)

- RN 173739-11-2 CAPLUS
- CN 1-Naphthalenol, 2-[(4-butyl-1-piperazinyl)methyl]-4-[(6-chloro-2-methoxy-9-acridinyl)amino]-5,6,7,8-tetrahydro- (CA INDEX NAME)

- RN 173739-12-3 CAPLUS
- CN 1-Naphthalenol, 4-[(6-chloro-2-methoxy-9-acridinyl)amino]-5,6,7,8tetrahydro-2-[[4-(2-methylpropyl)-1-piperazinyl]methyl]- (CA INDEX NAME)

- RN 173739-13-4 CAPLUS
- CN 1-Naphthalenol, 4-[(6-chloro-2-methoxy-9-acridinyl)amino]-5,6,7,8-tetrahydro-2-[[4-(1-methylpropyl)-1-piperazinyl]methyl]- (CA INDEX NAME)

- RN 173739-14-5 CAPLUS
- CN 1-Naphthalenol, 4-[(6-chloro-2-methoxy-9-acridinyl)amino]-5,6,7,8tetrahydro-2-[(4-pentyl-1-piperazinyl)methyl]- (CA INDEX NAME)

- RN 173739-15-6 CAPLUS
- CN 1-Naphthaleno1, 4-[(6-chloro-2-methoxy-9-acridiny1)amino]-5,6,7,8tetrahydro-2-[[4-(3-methylbuty1)-1-piperaziny1]methyl]- (CA INDEX NAME)

OH
$$CH_2-CH_2-CHMe_2$$
 $CH_2-CH_2-CHMe_2$ $CH_2-CH_2-CHMe_2$ $CH_2-CH_2-CHMe_2$ $CH_2-CH_2-CHMe_2$ $CH_2-CH_2-CHMe_2$

OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

L4 ANSWER 14 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1995:881293 CAPLUS DOCUMENT NUMBER: 123:286080

ORIGINAL REFERENCE NO.: 123:51271a,51274a

Preparation of

TITLE:

α-(mercaptoalkyl)-1-piperazineethanamines as inhibitors of farnesvl-protein transferase

INVENTOR(S):

Graham, Samuel L.; Williams, Theresa M. Merck and Co., Inc., USA

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 156 pp. CODEN: PIXXD2 Patent

DOCUMENT TYPE: LANGUAGE:

English FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.					KIND		DATE		APPLICATION NO.						DATE			
WO								WO 1994-US5634										
	₩:	ΑU,	BB,	BG,	BR,	BY,	CA,	CN,	CZ,	FI,	GE,	HU,	JP,	KG,	KR,	KZ,	LK,	
		LV,	MD,	MG,	MN,	MW,	NO,	NZ,	PL,	RO,	RU,	SD,	SI,	SK,	TJ,	TT,	UA,	
		US,	UZ															
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG			
CA	CA 2165176				A1 19950105				CA 1994-2165176						19940519			
AU	AU 9470412			A		19950117 AU 1994-70412							19940519					
AU	AU 675145				B2 19970123													
EP	EP 703905			A1 19960403				EP 1994-919174					19940519					
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LI,	LU,	NL,	PT,	SE	
JP	0950	0109			T		1997	0107		JP 1	994-	5028	10		1	9940	519	
ZA 9404326				A	A 19951214				ZA 1994-4326					19940617				
US	5736	539			A		1998	0407		US 1	995-	5498	29		1	9951	116	
PRIORITY APPLN. INFO.:									US 1	993-	8002	8	- 1	A 1	9930	618		
										US 1	994-	2375	86		A 1	9940	511	
										WO 1	994-	JS56	34	1	v 1	9940	519	

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OTHER SOURCE(S): MARPAT 123:286080

GI

Compds. which inhibit farnesyl-protein transferase (FTase) and the

farnesylation of the oncogene protein Ras were disclosed. More narrowly defined claimed compds. are $\alpha-(\text{mercaptomethyl})-1-$ piperazineethanamines I (Y = Ph, aryl, furanyl, etc.; Rl-R4 = H, alkyl,

substituent, etc.). The invention is further directed to chemotherapeutic compns. containing the compds of this invention and methods for inhibiting farnesyl-protein transferase and the farnesylation of the oncogene protein Ras.

T 169449-55-2P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of α -(mercaptoalkyl)-1-piperazineethanamines farnesyl-protein transferase inhibitors)

RN 169449-55-2 CAPLUS

1-Piperazinepropanethiol, β -amino-2-(2-methoxyethyl)-4-[(5,6,7,8-tetrahydro-1-naphthalenyl)carbonyl]-, [S-(R*,S*)]-, bis(trifluoroacetate) (salt) (9CI) (CA INDEX NAME)

CM 1

CN

CRN 169449-54-1 CMF C21 H33 N3 O2 S

Absolute stereochemistry.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

OS.CITING REF COUNT: 22 THERE ARE 22 CAPLUS RECORDS THAT CITE THIS

RECORD (22 CITINGS)

REFERENCE COUNT: 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

<12/04/2007>

Erich Leese

L4 ANSWER 15 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN 1994:508695 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: 121:108695

ORIGINAL REFERENCE NO.: 121:19627a,19630a

TITLE: Syntheses of Mannich basic compounds of tetrahydronaphthol containing piperazine side chains

AUTHOR(S): Gao, Shouhai; Li, Fulin

Inst. Pharmacol. Toxicol., Acad. Mil. Med. Sci., CORPORATE SOURCE:

Beijing, 100850, Peop. Rep. China

SOURCE: Zhongguo Yaowu Huaxue Zazhi (1993), 3(3), 175-8

CODEN: ZYHZEF; ISSN: 1005-0108

DOCUMENT TYPE: Journal LANGUAGE: Chinese GT

ΔR Title compds. I (R = Me, Et, Pr, Me2CH, Bu, iso-Bu, EtCHMe, pentyl, isopentyl) were prepared starting from 1-naphthol. I (R = Bu, EtCHMe,

isopentyl) showed antimalarial activity comparable to that of chloroquine.

156893-82-2P 156893-83-3P 156893-84-4P 156893-85-5P 156893-86-6P 156893-87-7P

156893-88-8P 156893-89-9P 156893-90-2P 156893-91-3P 156893-92-4P 156893-93-5P

156893-94-6P 156893-95-7P 156893-96-8P 156893-97-9P 156893-98-0P 156893-99-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation and antimalarial activity of)

RN 156893-82-2 CAPLUS

CN 1-Naphthalenol, 4-[(7-chloro-4-quinolinyl)amino]-5,6,7,8-tetrahydro-2-[(4methyl-1-piperazinyl)methyl]- (CA INDEX NAME)

RN 156893-83-3 CAPLUS

CN 1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-2-[(4-ethy1-1-piperaziny1)methy1]-5,6,7,8-tetrahydro- (CA INDEX NAME)

RN 156893-84-4 CAPLUS

CN 1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[(4-propy1-1-piperaziny1)methyl]- (CA INDEX NAME)

RN 156893-85-5 CAPLUS

<12/04/2007>

Erich Leese

CN 1-Naphthaleno1, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(1-methylethyl)-1-piperazinyl]methyl]- (CA INDEX NAME)

- RN 156893-86-6 CAPLUS
- CN 1-Maphthalenol, 2-[(4-butyl-1-piperazinyl)methyl]-4-[(7-chloro-4-quinolinyl)amino]-5,6,7,8-tetrahydro- (CA INDEX NAME)

- RN 156893-87-7 CAPLUS
- CN 1-Maphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(2-methylpropy1)-1-piperaziny1]methyl]- (CA INDEX NAME)

- RN 156893-88-8 CAPLUS
- CN 1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(1-methylpropy1)-1-piperaziny1]methyl]- (CA INDEX NAME)

- RN 156893-89-9 CAPLUS
- CN 1-Naphthaleno1, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[(4-penty1-1-piperaziny1)methy1]- (CA INDEX NAME)

RN 156893-90-2 CAPLUS

CN 1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(3-methylbuty1)-1-piperaziny1]methyl]- (CA INDEX NAME)

RN 156893-91-3 CAPLUS

CN 1-Naphthalenol, 4-1(7-chloro-4-quinolinyl)amino]-5,6,7,8-tetrahydro-2-1(4-methyl-1-piperazinyl)methyl]-, phosphate (1:3) (salt) (9C1) (CA INDEX NAME)

CM

CRN 156893-82-2

CMF C25 H29 C1 N4 O

CM 2

CRN 7664-38-2

CMF H3 O4 P

RN 156893-92-4 CAPLUS

CN 1-Naphthalenol, 4-[(7-chloro-4-quinolinyl)amino]-2-[(4-ethyl-1-piperazinyl)methyl]-5,6,7,8-tetrahydro-, phosphate (1:3) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 156893-83-3 CMF C26 H31 C1 N4 O

CM 2

CRN 7664-38-2 CMF H3 O4 P

CN

RN 156893-93-5 CAPLUS

1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[(4-propy1-1-piperaziny1)methy1]-, phosphate (1:3) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 156893-84-4 CMF C27 H33 C1 N4 O

<12/04/2007>

Erich Leese

CM 2

CRN 7664-38-2 CMF H3 O4 P

RN 156893-94-6 CAPLUS

CN 1-Naphthaleno1, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(1-methylethyl)-1-piperaziny1]methyl]-, phosphate (1:3) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 156893-85-5 CMF C27 H33 C1 N4 O

CM 2

CRN 7664-38-2 CMF H3 O4 P

о но- р- он он

RN 156893-95-7 CAPLUS
CN 1-Naphthalenol, 2-[(4-butyl-1-piperazinyl)methyl]-4-[(7-chloro-4-quinolinyl)amino]-5,6,7,8-tetrahydro-, phosphate (1:3) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 156893-86-6 CMF C28 H35 C1 N4 O

CM 2

CRN 7664-38-2 CMF H3 O4 P

RN 156893-96-8 CAPLUS

1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(2-methylpropy1)-1-piperaziny1]methyl]-, phosphate (1:3) (salt) (9CI) (CA

INDEX NAME)

CM 1

CRN 156893-87-7 CMF C28 H35 C1 N4 O

CM 2

CRN 7664-38-2 CMF H3 O4 P

CN

RN 156893-97-9 CAPLUS

1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(1-methylpropy1)-1-piperaziny1]methyl]-, phosphate (1:3) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 156893-88-8 CMF C28 H35 C1 N4 O

CM 2

CRN 7664-38-2 CMF H3 O4 P

RN 156893-98-0 CAPLUS

CN 1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,5,7,8-tetrahydro-2-[(4-penty1-1-piperaziny1)methy1]-, phosphate (1:3) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 156893-89-9 CMF C29 H37 C1 N4 O

CM 2

CRN 7664-38-2 CMF H3 O4 P

RN 156893-99-1 CAPLUS

CN 1-Naphthalenol, 4-[(7-chloro-4-quinoliny1)amino]-5,6,7,8-tetrahydro-2-[[4-(3-methylbuty1)-1-piperaziny1]methy1]-, phosphate (1:3) (salt) (9CI) (CA INDEX NAME)

CM 1

CRN 156893-90-2 CMF C29 H37 C1 N4 O

CM 2

CRN 7664-38-2 CMF H3 O4 P

CMF H3 04 P

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ANSWER 16 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                         1958:40588 CAPLUS
DOCUMENT NUMBER:
                         52:40588
ORIGINAL REFERENCE NO.: 52:7310h-i,7311a-i,7312a-e
                         Oxytocic activity of basic (aminomethyl) derivatives
TITLE:
                         of phenols and related compounds
AUTHOR(S):
                         Cohen, A.; Hall, R. A.; Heath-Brown, B.; Parkes, M.
                         W.; Rees, A. H.
CORPORATE SOURCE:
                         Roche Prods. Ltd., Welwyn Garden City, UK
SOURCE:
                         British Journal of Pharmacology and Chemotherapy
                         (1957), 12, 194-208
                         CODEN: BJPCAL; ISSN: 0366-0826
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         Unavailable
   The appropriate phenol, base, and formalin by the Mannich reaction gave
AB
     the following 2-naphthols [substituent, b.p./mm. or m.p. of base, other
     consts. given for base, m.p. of salts (HC1 = hydrochloride, T = acid
     tartrate, M = acid maleate)]: 1-(4-ethylpiperidinomethyl), 113°;
     1-(2-methylpiperidinomethyl), 94-6°; 1-(4-methylpiperidinomethyl),
     131.5-3.5°; 3-piperidino-methyl-5,6,7,8-tetrahydro (I),
     77-8°, HCl, 197-8°; 1-(2,4-dimethylpiperidinomethyl),
     71-3.5°: 1-(3-ethoxycarbonylpiperidinomethyl), -, HCl, 100°:
     1-(3-hydroxymethylpiperidinomethyl), -, M, 157-8°;
     1-(4-ethoxycarbonylpiperidinomethyl), -, HCl, 99-101°;
     3-(2-methylpiperidinomethyl)-5,6,7,8-tetrahydro, 120°/3 +
     10-5, n20D 1.552, T, 60-70°;
     3-(3-ethoxycarbonylpiperidinomethyl)-5,6,7,8-tetrahydro, 180°/0.3,
     HCl, 100°, T, 75-80°; 1-(3-methylpiperidinomethyl), -, M,
     157-8°; 1-(2-methyl-5-ethyl-piperidinomethyl), -, M, 70°;
     1-piperidinomethyl-3-ethoxycarbonyl, 106-8°, M, 121-3°;
     1-(α-piperidinoethyl), -, T, 125°. The following
     4,5-dimethylphenols: 2-(2-methylpiperidinomethyl) (II), -, HCl,
     190-2°, M, 134-6°; 2-(3-ethoxycarbonylpiperidinomethyl),
     116°/10-4, n20D 1.525; 2-(2,4-dimethylpiperidinomethyl),
     147°/0.5, n20D 1.527, HC1, 180-2°;
     2-(4-ethylpiperidinomethyl), 28-30°, HCl, 162-4°;
     2-(4-methylpiperidinomethyl), 44-6° HCl, 180-2°;
     2-(4-ethoxycarbonylpiperidinomethyl), 152°/5 + 10-5, n20D
     1.522, HCl, 164-6°; 2-(4-hydroxymethytpiperidinomethyl),
     75-6°, HCl, 180-2°; 2-(3-methylpiperidinomethyl),
     52-4°, -: 2-(2-methyl-5-ethylpiperidinomethyl), 80-1°,-;
     d-2-(2-methylpiperidinomethyl), 121°/0.3, n20D 1.534, [α]20D
     47.1° (c 0.98, benzene), -; 1-isomer, 112°/0.14, n20D 1.534,
     [\alpha]20D -51.4^{\circ} (c 1.33, benzene), -;
     2-hexamethyleniminomethyl, 52°, HCl, 174°;
     2-(N-ethyl-N-isopropylaminomethyl), 90°/0.15, n20D 1.516, HCl,
     201°; 2-isopropylaminomethyl, 75°, HCl, 137°;
     2-diethylaminomethyl, HCl, 190-2°; 2-morpholinomethyl.
     129°/0.4, HCl, 198°; 2-(2-ethylpiperidinomethyl), .apprx.
     39°, HCl, 174-6°; 2-(5-ethoxycarbonyl-2-
     methylpiperidinomethyl), -, HCl, 189°;
     2-(3-methylmorpholinomethyl), 59-61°, HCl, 165-6°, M, 162-4°; 2-diallylaminomethyl, 120°/0.1, HCl, 136-7°;
     2-dimethylaminomethyl, 80-1°, -; 2-pyrrolidinomethyl,
     130°/0.1, n20D 1.538, HCl, 149°. The following phenols:
     2-piperidinomethy1-3,5-dimethy1, -, M, 121-2°;
     6-piperidinomethy1-2,3-dimethy1, 128-30°/0.5, n20D 1.537, HC1,
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220-1.5°; 4-piperidinomethy1-2,5-dimethy1, -, HC1, 226-7°;
4-piperidinomethyl-2,6-dimethyl, -, M, 135-6°;
2-piperidinomethyl-4,6-dimethyl, -, M, 90°;
2-piperidinomethyl-3,4,6-trimethyl, -, HCl, 228-30°;
2-piperidinomethyl-4-methyl, -, HCl, 198°;
2-piperidinomethyl-5-methyl, -, HCl, 166-8°;
2-piperidinomethyl-4-chloro, -, HCl, 231°,
2-piperidinomethyl-4-chloro-5-methyl, -, HCl, 207°;
2-piperidinomethyl-4-ethyl-5-methyl, 120°/0.1, n20D 1.534, HCl,
160-2°; 2-piperidinomethyl-3,4,5-trimethyl, 102-3°, HCl,
211°; 2-(2-methylpiperidinomethyl)-4-ethyl-5-methyl,
143°/0.5, n20D 1.532, HCl, 143-5°;
2-piperidinomethyl-4,5-dimethoxy, 119°/5 + 10-5, HCl,
170-2°; 2-piperidinomethyl-4,5-diethyl, 136-8°/0.1, HCl,
178°; 2-piperidinomethyl-5-methyl-propyl, 132°/0.1, n20D
1.531, -; 2-piperidinomethyl-5-ethyl-4-methyl, 117°/0.1, HCl,
154°; 2-piperidinomethyl-4-propyl, 141°/0.75, n20D 1.528,
HCl, 178-80°; 2-(2-methylpiperidinomethyl)-5-ethyl-4-methyl,
126°/0.3, n20D 1.531, -; 2-piperidinomethyl-4-cyclohexyl,
59-60°, -; 1-2-(2-methylpiperidinomethyl)-4-ethyl-5-methyl,
126-8°/0.19, n20D 1.530, [α]20D -45.7° (c1.25,
benzene), -; d-isomer, 126-8°/0.19, n20D 1.530, [a]20D
44.4° (c 1.31, benzene), -:
2-piperidinomethyl-4-isopropyl-5-methyl, 122°/0.25, n20D 1.531, -.
The following 5-hydroxyindans: 6-piperidinomethyl, 125-6°/0.22,
n20D 1.549, HC1, 206-8°, M, 118°;
6-(2-methylpiperidinomethyl), 35-7°, HCl, 173-5°, M,
152-4°; 6-morpholinomethyl, 41-4°, M, 133°;
6-(3-methylmorpholinomethyl), 58-60°, HCl, 193-5°, M,
153°; 1-6-(2-methylpiperidinomethyl), 133-4°/0.1, n20D
1.549, [\alpha]20D -47.2° (c 0.68, benzene), M, 147-9° [[\alpha]20D -9.9° (c 1.7, water)]; d-isomer,136-8°/0.12,
n20D 1.549, [α]20D 44.9° (c 1.20, benzene), M, 144-7°
[[\alpha]20D 7.0° (c 1.63, H2O)]. The following compds.:
3-hydroxy-4-(piperidinomethyl)quinoline, m. 95°;
6-hydroxy-5-(piperidinomethyl)quinoline-HCl, m. 214°; and
3-(β-piperidinoethyl)indole-HCl, m. 222.5-4.5°.
1-Bromo-5,6,7,8-tetrahydro-2-naphthol in a Mannich reaction gave
1-bromo-3-piperidinomethyl-5,6,7,8-tetrahydro-2-naphthol from which Br was
eliminated by hydrogenation in HOAc with PdBaSO4 in the presence of KOAc
to give I. Also 2-hydroxy-5,6,7,8-tetrahydro-3-naphthoic ester, converted
to the piperidide, m. 202-4°, on reduction with LiAlH4 gave I.
2-Hydroxy-3-naphthopiperidide, prisms, m. 229-30° (MeOH), prepared
from 3-ethoxycarbonyl-2-naphthol, on reduction with LiAlH4 gave
3-piperidinomethyl-2-naphthol, m. 159-60°; HCl salt, m.
217.5-19.5°. 2-Bromo-4.5-dimethyl-phenol by a Mannich reaction
gave 2-bromo-4.5-dimethyl-6-piperidinomethylphenol, m. 93-5°,
debrominated as above to 2-piperidinomethyl-3,4-dimethylphenol, b0.18
120-2°; HCl salt, m. 168-70°. Salicylaldehyde and
piperidine hydrogenated with Pd-C catalyst gave 2-piperidinomethylphenol,
b0.25 100°, n20D 1.537; HCl salt, m. 160-2°. The Kindler-Willgerodt reaction with 2-benzyloxy-4,5-dimethylacetophenone gave
a substituted phenylacetothiomorpholide, m. 129°, which
desulfurized with Raney Ni gave 1-β-[(2-benzyl-oxy-4,5-
dimethylphenyl)ethyl]morpholine; picrate, m. 178-8.5°.
Hydrogenation of the crude base HCl salt with Pd-C gave
2-(β-morpholinoethyl)-4,5-dimethylphenol-HCl, m. 238-9°.
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Similarly, from the phenylacetothiopiperidide was obtained 1-B-((2-benzyloxy-4,5-dimethylphenyl)ethyllpiperidine-HCl, m. 180-1°, hydrogenated to 2-(β-piperidinoethyl)-4,5dimethylphenol-HCl, m. 193-5°. 2-Amino-4,5-dimethylphenol, 1,5-dibromopentane, and K2CO3 in boiling BuOH gave 2-piperidino-4,5-dimethylphenol, b0.1 95-7°, n20D 1.539. II was converted to the acetoxy derivative, b0.05 118°, n20D 1.527 and to the benzovloxy derivative, m. 77-8°, by treating 20 hrs. at 20° with the corresponding chloride in dry pyridine. A mixture of 5-methoxyindan-6-aldehyde and α-pipecoline hydrogenated over Pd-C gave 5-methoxy-6-(2-methylpiperidinomethyl)indan, b0.05 129-31°, n20D 1.543. These compds. were tested for oxytocic activity both in vivo and in vitro and some were found to exceed ergometrine in activity. Highest activity occurred with 2-piperidinomethyl derivs. of phenols, among which maximum potency was conferred by substitution at both the 4 and 5 positions by Me or Et or by linkage of these positions to form an indan derivative In all series, piperidinomethyl derivs. were more active than those formed with other bases and methylation in the position α to the N atom augmented the activity of both piperidine and morpholine derivs. Among 2-methylpiperidinomethyl phenols, the 1- was more active than the d-form. Acylation or alkylation of the phenolic HO group did not affect activity. The oxytocic activity was specific, the compds. being less effective upon other forms of smooth muscle. Effects upon blood pressure and respiration of a central nature were observed. 1071701-96-6P

RI: SPN (Synthetic preparation); PRP (Properties); PREP (Preparation) (Oxytocic activity of basic (aminomethyl) derivatives of phenols and related compounds) 1071701-96-6 (APLUS

RN 1071701-96-6 CAPLUS
CAPLUS
Butanedioic acid, 2,3-dihydroxy-, 1-[3-[3-(ethoxycarbonyl)-1-piperidinyl]methyl]-5,6,7,8-tetrahydro-2-naphthalenyl] ester (CA INDEX NAME)

IT 860440-00-2P, 2-Naphthol, 5,6,7,8-tetrahydro-3-piperidinomethyl-860440-02-4P, 2-Naphthol, 5,6,7,8-tetrahydro-3-piperidinomethyl-, hydrochloride RL: PREP (Preparation)

(preparation of)

RN 860440-00-2 CAPLUS

CN 2-Naphthalenol, 5,6,7,8-tetrahydro-3-(1-piperidinylmethyl)- (CA INDEX NAME)

RN 860440-02-4 CAPLUS

CN 2-Naphthalenol, 5,6,7,8-tetrahydro-3-(1-piperidinylmethyl)-, hydrochloride (1:1) (CA INDEX NAME)

HCl

OS.CITING REF COUNT: 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS)

L4 ANSWER 17 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1957:101470 CAPLUS

DOCUMENT NUMBER: 51:101470

ORIGINAL REFERENCE NO.: 51:18343i,18344a-c

TITLE: Pharmacological research on synthetic uterotonics. II.

Substituted N-benzylpiperidines and

3,4-dimethoxybenzylamines
AUTHOR(S): Votava, Z.; Podvalova, I.

CORPORATE SOURCE: Research Inst. Pharmacy and Biochemistry, Prague

SOURCE: Chekhoslovatskaya Fiziologiya (1954), 3, 426-31

CODEN: CHFIAK; ISSN: 0031-9309

DOCUMENT TYPE: Journal LANGUAGE: English

AB cf. C.A. 50, 8900c. Tests were carried out for pharmacol. properties of the following N-benzylpiperidine derivs.: 3,4-tetramethylene; 2-methoxy; 3-methoxy; 4-methoxy; 2,3-dimethoxy; 2,6-dimethoxy; 2,4-dimethoxy; 2,5-dimethoxy; 2-hydroxy-5-methoxy; 2,6-dimethoxy; 3,4-dimethoxy;

1-methyl-3,4-dimethoxy; 3,4-methylenedioxy; 3,4-ethylenedioxy; 3-methoxy-4-hydroxy; 3,5-dimethoxy; 2,3,4-trimethoxy; 2,4,5-trimethoxy;

3,4,5-trimethoxy; and 4-hydroxy-3,5-dimethoxy; the following N,N-disubstituted derivs. of 3,4-dimethoxybenzylamine: di-Me; di-Et; di-Pu; di-Bu; and diallyl and the N-(3,4-dimethoxybenzyl) derivs. of: pyrrolidine; piperidine; 2-methylpiperidine; 2,6-dimethylpiperidine;

hexamethylenimine; 1-[1-(3,4-dimethoxyphenyl)ethyl]piperidine; and 1-(3-indolylmethyl)-2-methylpiperidine. In all substances, the uterotonic action was studied on in situ expts. in rabbits, the effect on the blood pressure in rabbits, and the toxicity in mice. The substances were always

pressure in rabbits, and the toxicity in mice. The substances were always administered intravenously. A regularity was determined between the chemical structure and the uterotonic effect of the substance. 860227-77-6, Piperidine,

1-[(5,6,7,8-tetrahydro-2-naphthyl)methyl]-

(pharmacology of) RN 860227-77-6 CAPLUS

CN Piperidine, 1-[(5,6,7,8-tetrahydro-2-naphthalenyl)methyl]- (CA INDEX NAME)

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)

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L4 ANSWER 18 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                       1956:48646 CAPLUS
DOCUMENT NUMBER:
                        50:48646
ORIGINAL REFERENCE NO.: 50:9354g-i,9355a-q
                        ar-2-Tetralol derivatives
AUTHOR(S):
                        Hull, Robert L.
SOURCE:
                        Journal of the American Chemical Society (1955), 77,
                        6376-9
                        CODEN: JACSAT; ISSN: 0002-7863
DOCUMENT TYPE:
                        Journal
LANGUAGE:
                        Unavailable
OTHER SOURCE(S):
                        CASREACT 50:48646
GI For diagram(s), see printed CA Issue.
    1-(Piperidinomethyl)-2-naphthol (72.4 g.) in 200 cc. glacial AcOH
    hydrogenated 20 h. at 60° and 50 lb. over 3.0 g. 5% Pd-C, the mixture
     filtered into 1 1. ice water, the precipitate filtered off, shaken with 300 cc.
     Et20 and 300 cc. H20, the Et20 extract dried, evaporated, and the solid residue
     recrystd. from ligroine (b. 60-70°) gave 29.4 g.
     5,-6,7,8-tetrahydro-1-methyl-2-naphthol (I), colorless needles, m.
     113-14°. I (16.2 g.) and 8.5 g. piperidine in 50 cc. EtOH treated
     with 8.2 g. 36-8% agueous CH2O, the mixture allowed to stand overnight, cooled
     in ice, filtered, and the filter cake washed with cold EtOH vielded 18.1
    q. 3-(piperidinomethyl)-derivative of I, m. 57-9°; the mother liquor
     concentrated gave an addnl. 6.8 g. material, m. 57-9°; anal. sample, m.
     60.5-1.5° (from EtOH). I (32.4 g.) in 200 cc. CC14 treated
     dropwise during 15 min. with 27.0 g. SO2Cl2, the mixture washed with 300 cc.
     H2O, 300 cc. 5% aqueous NaHCO3, and 300 cc. H2O, dried, evaporated on the steam
     bath, the residual oil distilled and the fraction b0.5 90-115°, which
    solidified, recrystd. from 75 cc. 70% EtOH gave 21.0 g. 3-Cl derivative of I,
    colorless needles, m. 57-8° (from EtOH). Br (32 g.) in 50 cc. CC4
    added dropwise with stirring to 32.4 g. I in 150 cc. CC14, the solution
    stirred 0.5 h., washed with 300 cc. H2O, 300 cc. 5% aqueous NaHCO3, and 500
    cc. H2O, dried, evaporated, and the solid residue recrystd. from 70% EtOH gave
    36.5 g. 3-Br derivative of I, colorless needles, m. 69-70°.
     5,6,7,8-Tetra-hydro-3-Pr 2-naphthol (II) treated with Br in CC14 yielded
     53% 1-Br derivative of II, m. 64.5-5.5° (from 70% EtOH). The
     appropriate ar-2-tetralol (0.05 mol) in 25 cc. absolute EtOH added to 1.15 q.
    Na in 20 cc. absolute EtOH, the mixture treated with HOCH2CH(OH)CH2Cl or
    HOCH2CMe(OH)CH2Cl, refluxed 3 h., filtered, the filtrate evaporated, and the
    residue recrystd, or distilled gave the corresponding III (R. X. Y. % vield,
     and m.p. given): H, Me, H, 42, 109-10°; Me, Me, H, 34,
    91.5-2.5°; Me, H, H, 70, 80-1°; H, Br, H, 31, 120-1°;
    H, Br, Br, 42, 104.5-5.5°; H, Me, Br, 29, 85.5-6.5°; H, Me,
     Cl, 24, 81-2°; Me, Me, Br, 51, 102.5-3.5°; H, H, CH2CH:CH2,
     21, 66-7°; H, H, Pr, 47, 88-9°; Me, H, CH2CH:CH2, 44, -
    (b0.5 175-80°); Me, Br, Br, 23, 81-2°, H, Br, Pr, 51,
     86-7°; Me, Br, Pr, 23, 80-1°.
     1,3-Dibromo-5,6,7,8-tetrahydro-2-naphthyl acetate (10.4 q.) added to 2.8
     q. NaOH in 40 cc. 70% EtOH, the mixture refluxed 1 h., treated with 0.040
    mol of the appropriate glycerol monohydrin, refluxed 3 h., evaporated in vacuo
     at 50°, the gummy residue extracted with 100 cc. hot C6H6, the extract
     evaporated, and the residue recrystd. gave the III (X and Y = Br). I (34.5
     g.) and 20 cc. concentrated H2SO4 heated 0.5 h. on the steam bath, the deep red
     solution diluted with 150 cc. H2O, cooled in ice, treated with stirring with 14
     cc. concentrated HNO3, the mixture heated 10 min. on the steam bath, diluted
with an
     equal volume of H2O, cooled in ice, and the yellow precipitate filtered, washed
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with H2O, and recrystd. from EtOH gave 26.4 q. 3-nitro derivative (IV) of I, yellow needles, m. 118-19°. II nitrated in the same manner yielded 60% 1-nitro derivative (V) of II, yellow needles, m. 104-5° (from EtOH). IV (10.4 g.) in 200 cc. absolute EtOH hydrogenated 10 min. at room temperature and 50 lb. pressure over 0.1 q. PtO2, the mixture filtered, the filtrate diluted with 4 vols. H2O, and the precipitate filtered off, dried (8.5 q.), and recrystd. from ligroine gave the 3-amino derivative (VI) of I, m. 144-5°. V hydrogenated in the same manner vielded 71% 1-amino derivative (VII) of II, m. 95-6° (from aqueous EtOH). VI (8.1 q.) and 25 cc. 98% HCO2H refluxed 1 h., the excess HCO2H and H2O distilled off, the residue heated 4 h. at 140-50°, the cooled solid extracted with two 50-cc. portions of Me3CCH2CHMe2, the extract cooled in ice, and the white crystalline deposit (5.7 g.) recrystd. from EtOH gave 5,6,7,8-tetrahydro-9-methylnaphth[2,3]oxazole (VIII), colorless crystals, m. 94-5°. VII gave similarly 60% 6,7,8,9-tetrahydro-4-propylnaphth[1,2]oxazole (IX), colorless oil, b0.1 99-101°. NH2OH.HC1 (1.3 g.) and 3.4 g. VIII added to 0.8 g. NaOH in 25 cc. H2O and 30 cc. EtOH, the mixture refluxed 0.5 h., diluted with an equal volume of H2O, cooled in ice, and the cream-colored solid deposit filtered, dried (2.6 g.), and recrystd, from EtOH-C6H6 gave the 2-NH2 derivative of VIII.H20, m. 159-60°. IX gave similarly 64% 2-NH2 derivative of IX, m. 174-5° (from ligroine).

II 412014-25-6P, Piperidine,
1-[(5,6,7)8-tetrahydro-3-hydroxy-4-methyl-2-naphthyl)-methyl]RL: PREP (Preparation of)
(preparation of)

RN 412014-25-6 CAPLUS

CN 2-Naphthalenol, 5,6,7,8-tetrahydro-1-methyl-3-(1-piperidinylmethyl)- (CA INDEX NAME)

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD
(1 CITINGS)

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L4 ANSWER 19 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER:
                        1956:40401 CAPLUS
DOCUMENT NUMBER:
                         50:40401
ORIGINAL REFERENCE NO.: 50:7803c-f
                        Chloromethylation of tetralin
TITLE:
AUTHOR(S):
                        Vanags, G.; Gudriniece, E.
SOURCE:
                         Latvijas PSR Zinatnu Akademijas Vestis (1955), (No. 5
                         (Whole No. 94)), 119-24
                         CODEN: LZAVAL; ISSN: 0132-6422
DOCUMENT TYPE:
                        Journal
LANGUAGE:
                        Russian
     Tetralin (66 mg.), 28 g. (CH2O)n, 65 ml. glacial AcOH, 33 g. crystalline H3PO4,
     and 91 ml. concentration HCl at 85-90° stirred 4 hrs. gave 66%
     1,2,3,4-tetrahydro-6-chloromethylnaphthalene (I). With excess II, 10%
     5,8-bis(chloromethyl)-1,2,3,4-tetrahydronaphthalene was obtained in addition
     to I. The 6-piperidinomethyl analog (II of I) was prepared by treating I in
     Et20 with piperidine at room temperature II decomposed on distillation
Bubbling dry HCl
     through II in Et20 gave II.HCl, very hygroscopic. II picrate, m.
     150°. 1-(1,2,3,4-Tetrahydro-6-naphthylmethyl)pyridinium chloride,
     m. 115°, was prepared (88.5% yield) from 7.2 g. I, 20 ml. absolute Et20,
     and dry pyridine. H2NC(SR):NH.HCl (R =
     1,2,3,4-tetrahydro-6-naphthylmethyl), m. 212°, was prepared (96%
     yield) by heating 7.2 g. I with 6 g. thiourea. RCO2H was prepared (42%
     yield) refluxing crude I with KCN in H2O, and hydrolyzing the nitrile with
     aqueous NaOH; the hydrolysis was aided, and formation of resinous products was
     minimized by adding small amts. of 3% H2O2 at intervals. RCONHPh, m.
     112°, was obtained by method similar to that described (C.A. 50,
     271f).
    860227-77-6, Piperidine,
     1-[(5,6,7,8-tetrahydro-2-naphthyl)methyl]-
        (and derivs.)
     860227-77-6 CAPLUS
RN
CM
     Piperidine, 1-[(5,6,7,8-tetrahydro-2-naphthalenyl)methyl]- (CA INDEX
    NAME)
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IIT 857435-57-5P, Pyridinium,
1-[(5,6,7,8-tetrahydro-2-naphthyl)methyl]-, chloride
RL: PREP (Preparation)
 (preparation of)

RN 857435-57-5 CAPLUS

CN Pyridinium, 1-((5,6,7,8-tetrahydro-2-naphthalenyl)methyl]-, chloride (1:1)
(CA INDEX NAME)

• c1-

L4 ANSWER 20 OF 20 CAPLUS COPYRIGHT 2009 ACS on STN

ACCESSION NUMBER: 1950:10095 CAPLUS

DOCUMENT NUMBER: 44:10095

ORIGINAL REFERENCE NO.: 44:1979i,1980a-b

TITLE: Piperidylmethyl compounds with oxytocic action AUTHOR(S): Schindler, O.; Voegtli, W.

SOURCE: Pharmaceutica Acta Helvetiae (1949), 24, 207-16

CODEN: PAHEAA; ISSN: 0031-6865

DOCUMENT TYPE: Journal Unavailable

LANGUAGE:

Procedures for preparing the following compds. are given:

2-(1-piperidylmethyl)-5,6,7,8-tetrahydronaphthalene,

2-(1-piperidylmethyl)-1-chlorocyclohexane,

1-(1-piperidylmethyl)cyclohexene, 2-(1-piperidylmethyl)-1-

chlorocyclopentane, and 1-(1-piperidylmethyl)cyclopentene. These compds. appear to have about 0.1 the activity of methylergobasine when tested on the uterus of the guinea pig. 22 references.

860227-77-6, Piperidine,

1-[(5,6,7,8-tetrahydro-2-naphthy1)methy1]-(and derivs.)

860227-77-6 CAPLUS RN

Piperidine, 1-[(5,6,7,8-tetrahydro-2-naphthalenyl)methyl]- (CA INDEX CN NAME)

=> d his

(FILE 'HOME' ENTERED AT 17:55:20 ON 23 NOV 2009)

FILE 'REGISTRY' ENTERED AT 17:55:27 ON 23 NOV 2009

L1 STRUCTURE UPLOADED

L2 0 S L1 SSS L3 190 S L1 FULL

FILE 'CAPLUS' ENTERED AT 17:56:37 ON 23 NOV 2009 L4 20 S L3 FULL

=> log y

 COST ÎN U.S. DOLLARS
 SINCE FILE ENTRY
 TOTAL ENTRY

 FULL ESTIMATED COST
 125.30
 311.88

 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
 SINCE FILE ENTRY ENTRY
 TOTAL ENTRY

 CA SUBSCRIBER PRICE
 -16.40
 -16.40

STN INTERNATIONAL LOGOFF AT 18:11:52 ON 23 NOV 2009